Distance Histogram Computation Based on Spatiotemporal Uniformity in Scientific Data

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Abstract The large amount of data generated by scientific applications introduces a great challenge in data management and storage. Many queries against scientific data are analytical in nature and require superlinear time to process. This paper focuses on an important query in scientific simulation data analysis - the Spatial Distance Histogram (SDH). The computation time of an SDH query using brute force method is quadratic. Furthermore, such types of queries are often executed continuously to analyze the simulation system over certain time periods. This increases the total computation time of the SDH. We propose an approximate but highly efficient algorithm to compute SDH over consecutive time periods with provable error bounds. The key idea of our algorithm is to derive statistical distribution of distances from the spatial and temporal characteristics of particles. Upon organizing the data into a Quad-

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tree based structure, the spatiotemporal characteristics of particles in each node of the tree are acquired to determine the particles spatial distribution as well as their temporal locality in consecutive time periods. We also report our efforts in implementing and optimizing the above algorithm in Graphics Processing Units (GPUs) as a means to further improve the efficiency. The accuracy and efficiency of the proposed algorithm is backed by mathematical analysis and results of extensive experiments using data generated from real simulation studies.

Keywords Scientific databases \cdot spatial distance histogram \cdot quad-tree \cdot density map \cdot spatiotemporal locality \cdot GPU

1 Introduction

The advancement of computer simulation systems and experimental devices has yielded large volume of scientific data. This imposes great strain on the data management software, in spite of effort made to deal with such large amount of data using database management systems (DBMS) [11,18,32]. But the traditional DB-MSs are built with business applications in mind and are not suitable for managing the scientific data. Therefore, there is a need to have another look at the design of the data management systems. Data in scientific databases is generally accessed through high-level analytical queries, which are much more complex to compute in comparison to simple aggregates. Many of these queries are composed of few frequently used analytical routines which usually take super-linear time to compute using brute-force methods. Hence, the scientific database systems need to be able to efficiently handle the computation of such analytical queries. This

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Fig. 1 A snapshot of simulated collagen fiber structure

paper presents our work related to such type of a query that is very important for the analysis of *molecular simulation* (MS) data.

Molecular (or particle) simulations are simulations of complex physical, chemical or biological structures done on computers. They are extensively used as a basic research tool in material sciences, astronomical physics, biophysics, biomedical sciences, etc. The core units of these simulations are natural particles (e.g., atoms, molecules, etc.) that interact with each other governed by classical forces. Researchers perform such simulations to analyze the behavior of natural systems under experimental framework based on principal theoretical models [14,31]. The number of particles involved in MSs is usually large, oftentimes counting millions. For example, Fig. 1 is a visualization of a section of a collagen fiber system and this section along contains more than 890,000 atoms. In addition to the large amount of particles, simulation datasets may consist of multiple snapshots (called *frames*) of the system's state captured at different time points. Each frame contains measurements (e.g., spatial coordinates, mass, velocity, charge) of all particles and big number (e.g., tens of thousands) of such frames are being stored during a typical simulation process.

In order to analyze the MS data, scientists compute complex quantities through which statistical properties of the data is shown. Often times, queries used in such analysis count more than one particle as basic unit: such a function involving all *m*-tuple subsets of the data is called an m-body correlation function. One such analytical query discussed in this paper, is the so called spatial distance histogram (SDH) [42]. An SDH is the histogram of distances between all pairs of particles in the system and it represents a discrete approximation of the continuous probability distribution of distances named Radial Distribution Function (RDF). Being one of the basic building blocks for a series of critical quantities (e.g., total pressure and energy) required to describe the physical systems, this type of query is very important in MS databases [14].

1.1 Problem Statement and Notations

The SDH problem can be formally described as follows: given the coordinates of N particles and a userdefined distance w, we need to compute the number of particle-to-particle distances falling into a series of ranges (named buckets) of width $w: [0, w), [w, 2w), \ldots$, [(l-1)w, lw]. Essentially, the SDH provides an ordered list of non-negative integers $H = (h_0, h_1, \ldots, h_{l-1})$, where each $h_i (0 \le i < l)$ is the number of distances falling into the bucket [iw, (i + 1)w). We also use H[i] to denote h_i in this paper. Clearly, the bucket width w (or total bucket number l) is the only parameter of this type of problem.

To capture the variations of system states over time, there is also the need to compute SDH for a large number of consecutive frames. We denote the count in bucket i at frame j as $H_j[i]$.

Notations used throughout this paper are listed in Table 1. If a notation is not shown in this table, it is used in a local context only.

 Table 1 Common notations and their definitions.

Symbol	Definition
N	total number of particles in data
l	total number of histogram buckets
w	width of histogram buckets
i	an index on histogram buckets
j	an index on frames
H[i]	counts in the i -th bucket of the SDH
M	number of cells in the density map of interest
n_A	number of particles in a cell A
r_A	count ratio of a cell A
α	probability bound in chi-square test for identi-
	fying uniform regions

1.2 Overview of Our Approach.

Previous work [9,42,17] has studied algorithms to compute SDHs. Proved to have low time complexity, such algorithms, however, are not always *practical* in that the running time can still be long under certain conditions.¹ In contrast to those, this paper presents a **highly efficient and practical** algorithm for processing SDH of large-scale MS data. Our algorithm shows efficiency that is orders of magnitude higher than existing solutions under a wide range of query parameters. On the other hand, the accuracy of query results is also higher in our algorithm. To achieve the desired efficiency and accuracy, the algorithm mainly takes advantage of the two types of uniformity widely present in MS

¹ Details can be found in Section 2.

data. To further improve the running time of the algorithm, we harness the computational power of modern massively parallel hardware by implementing and optimizing the proposed algorithm in Graphics Processing Unites (GPUs).

The first type of data uniformity used by the algorithm refers to the spatial distribution of data **points** (e.g., atoms) in MS datasets. We have observed that systems generated by molecular simulations often have regions in the simulation space where the data particles are uniformly distributed. Also, it is well known that parts of natural systems tend to spread out evenly in space due to the existence of inter-particle forces and/or chemical bonds [3,5]. Because of this, there are many localized regions in the simulation space in which the particles are uniformly distributed.² For instance, Fig. 1 shows that the solvent molecules, represented by red dots, are uniformly distributed since a body of water is basically incompressible. Because of this uniformity, we treat such regions as single entities when computing SDH. As shown later in this paper, utilizing this type of uniformity does not introduce significant error to the accuracy of the algorithm. The main achievement via exploiting this property is that it makes the running time of the algorithm independent to the SDH bucket width w – such dependency (as discussed in Section 2) is the main drawback of existing algorithms. This technique is presented in more details in Section 4.

The second type of uniformity is about the significant **temporal similarity among neighboring frames**. We have observed that such similarity is actually reflected in the final results of the SDH obtained for neighboring frames. So, given two frames f_0 and f_1 , if we have already computed the SDH of f_0 , we can obtain the SDH of f_1 by dealing only with the regions that do not exhibit similarity between the two frames while ignoring regions that are similar. To take advantage of such similarities among frames, we design an *incremental algorithm* that can quickly compute SDH of a frame from the SDH of a base frame which was obtained using traditional single-frame algorithms. Section 5 has more details on this technique.

Finally, parallel processing is an obvious strategy to reduce computing time in our problem. As mentioned earlier, our algorithm takes advantage of the processing power of the GPUs. GPUs are popular, massively parallel systems used in many scientific applications. Because of their highly parallel structure, they are more effective than general purpose CPUs especially for algorithms like ours that can process large blocks of data in parallel. For us, they provide a low-cost and lowpower platform to improve efficiency as compared to computer clusters. However, the unique architecture of GPUs imposes interesting challenges to developing software that takes full advantage of the computing power of GPUs. In this paper, we develop several techniques to address such challenges. Such techniques are very different from those used for optimizing towards CPU-based systems and generate significant boosts in performance (and energy efficiency) as compared to straightforward GPU implementations. More technical details about the implementation of our algorithm to harness the GPU power can be found in Section 8.

1.3 Contributions and Paper Organization

We have implemented a composite algorithm combining the above ideas and tested it on real MS datasets. The experimental results clearly show the superiority of the proposed algorithm over previous solutions in both efficiency and accuracy. For example, with the proposed algorithm, we are able to compute 11 frames of a 8million-atom dataset in less than a second! In addition to a highly efficient and practical algorithm for SDH processing, we also believe that our success will open up new directions in the molecular simulation paradigm. First, our work builds a solid foundation for solving the more general and also difficult problem of multi-body (m-body) correlation function computation [38]. With a $O(N^m)$ complexity in nature, such problems can be addressed by taking advantage of the methodologies we propose in this paper. Second, the high efficiency of our algorithm enables on-the-fly MS data processing: data can be analyzed as they are generated. As a result, we can integrate our algorithm into the simulation software such that effective tuning of the simulation process becomes feasible.

The major technical contributions of our work presented in this paper are:

- Techniques to identify spatial uniformity within a frame and temporal uniformity among consecutive frames;
- An approximate algorithm to compute the SDH of large number of data frames by utilizing the above properties;
- Analytical and empirical evaluation of the above algorithm, especially rigorous analysis of the tradeoff between performance and guaranteed accuracy of the algorithm; and
- Implementation of the above algorithms in modern GPUs to boost performance, with a focus on the optimization of such implementations in a GPU programming environment.

² This does not make the data system-wise uniform. Otherwise, SDH computation becomes a trivial task.

The remainder of this paper is organized as follows: in Section 2 we give an overview of the work done in the field related to the SDH problem. Then, in Section 3 we introduce the main concepts and techniques utilized in our work. Sections 4 and 5 discuss the utilization of the spatio-temporal properties of the data to enhance the algorithm. In Section 6 we present an algorithm that combines both spatial uniformity and temporal locality properties. Then, in Section 7 the performance (running time and errors) of the proposed technique in utilizing the spatio-temporal property of the data is analyzed. In Section 8 we briefly look at the basic architecture of the GPUs and their programming paradigms and we modify our algorithm to map onto the GPU. Section 9 presents the results obtained through extensive experiments. Finally, we conclude this paper with Section 10 in which we also discuss our future work.

2 Comparison to Related Work

The brute-force method for SDH computation calculates the distances between all the pairs of particles in the MS system and distributes these distances into the relevant buckets of the histogram. This method requires quadratic time. Some of the popular software for analyzing the MS data, like GROMACS [23], still utilizes the brute-force method for SDH computation. But, the current state-of-the-art models for SDH computation involve methods that treat a cluster of particles as a single processing unit [42,17]. Space-partitioning trees (like kd-trees [17]) are often used to represent the system, each node of the tree representing one cluster. The main idea in such approach is to process all the particles in each node of the tree as a whole. This is an obvious improvement in terms of time over the bruteforce method which builds the histogram by computing particle-to-particle distances separately. A Density-Map based SDH algorithm (DM-SDH) using a quadtree data structure is presented in our previous work [42] It has been proved that the running time for DM-SDH is $\Theta(N^{\frac{3}{2}})$ for 2D data and $\Theta(N^{\frac{5}{3}})$ for 3D data. We will go over the main idea of DM-SDH in more detail later in this paper. Although the DM-SDH algorithm is an improvement over the brute-force method for SDH computation, it is still not a practical and efficient solution for the following reasons:

(1) The running time analysis of DM-SDH [42] is done considering the data size N as input (under a certain bucket width w). Considering the running time as a function of w, or the total number of buckets l in the SDH, one can see that the running time increases dramatically with w decreases (or l increases). As a result, when w is considerably small, the running time of the DM-SDH can even be greater than that of the brute-force method [42]!

(2) DM-SDH only addresses the SDH computation of a single frame. We mentioned earlier in this paper how important is the MS data analysis of the system over a period of time, i.e. multiple consecutive frames. In order for DM-SDH algorithm to achieve such computation over F consecutive frames, one needs to actually run the same algorithm F times. This is not quite acceptable, since usually F is on the order of tens of thousands.

An approximate SDH algorithm (ADM-SDH), with running time not related to the data size N was also introduced in [42]. But its running time is influenced by a guaranteed error bound as well as by the bucket size w. Like the DM-SDH, it too can only be applied to a single frame of the MS system. A thorough analysis of the performance of ADM-SDH is presented in a recent paper [21]. Under some assumptions, that paper also derives an error bound of ADM-SDH that is tighter than the one presented in [42]. We will briefly mention such findings in Section 7.1.2. To remedy the cons of these two aforementioned algorithms, we direct our current work in designing a new, improved algorithm with higher efficiency and accuracy. Furthermore, we are able to substantially decrease the running time of the algorithm by implementing and optimizing code in a GPU programming environment. The end result is a solution that is both practical and efficient, delivering very accurate results in a (almost) real-time manner. A shorter version of this paper can be found in [29].

It is important to note here that the SDH problem is often confused with the force/potentional fields computation in the MS process [6,19]. In the latter, the physical properties of a particle are determined by the forces applied to it by all other particles in the system. Different approximate algorithms have been introduced to tackle this problem by making use of the mathematical features of the formula that defines such forces. Even though the force/potentional fields computation has similar definition to the SDH problem, the algorithms used to solve such problem are not useful in computing the SDH. There is a detailed comparison of the two problems in [9]. Here, we will just note that the force field computation is for simulation of a system, while the SDH computation is for system analysis.

The problem of SDH computation over multiple consecutive frames is related to persistent data structures [10], which allow for various versions of the computation results to be maintained and updated over time for quick query processing. Such a structure can be extended into a multi-dimensional persistent tree (i.e., MP-tree) [40] for searching in spatio-temporal data. Efficient handling of spatio-temporal data has been motivated by location based services (such as GIS application). Building persistent index schemes on complex spatio-temporal data allows for a time efficient retrieval [30]. There has been a detailed survey of applications made by Kaplan [26] in which persistent data structures has been used to improve efficiency. Such structures are designed to resolve the I/O bottleneck problem. But, the multi-frame SDH problem involves heavy computation at each instance, overshadowing the I/O time. Thus, the techniques developed for persistent data can hardly be used for efficient multi-frame SDH computation.

Special hardwares fulfill the need of computation intensive tasks. In recent years, there is strong interest in improving the performance of such tasks using GPUs [24,28]. These devices were originally designed for processing graphics, but their computing power can be utilized for general purpose computing via software frameworks such as CUDA [34] and OpenCL [27]. A number of database operators are implemented on GPUs: relational join processing on GPUs is presented by He et al. [22] while a number of algorithms for other relational operators including selections, aggregations [16] as well as sorting [15] are presented by Govidaraju *et* al.. An overview of the GPU techniques is available in a detailed survey of general purpose computation on graphics hardware done by Owens et al. [37]. In this work, we leverage the computing power of the GPUs to achieve the goal of on-the-fly SDH computation with guaranteed accuracy.

3 Background

In this section, we introduce the main concepts and basic techniques presented in existing work [42] but will serve as a foundation for the proposed algorithm. To represent the simulation data space we use a conceptual data structure we call *density map* (DM). A DM divides the simulation space into a grid of equal sized cells (or regions). A cell is a square in 2D and a cube in 3D.³ To generate a density map of higher resolution, we divide each cell of the grid into four equally sized cells. We use a region quad-tree [36] to organize different density maps of the same data. Each cell of the DM is represented by a tree node, so a density map is essentially the collection of all nodes on **one level** of the tree. Each node of the tree contains the cell location (i.e., coordinates of corner points) as well as the



Fig. 2 Computing minimum (i.e., length of solid lines) and maximum distance (i.e., length of dashed lines) range between two cells

number of particles in it. We refer to such a tree as the Density-Map Tree (DM-tree).

The fundamental part of the DM-SDH algorithm is a procedure we call RESOLVETWOCELLS. This procedure takes two cells (e.g., A and B in Fig. 2) from a density map as an input and computes the minimum and maximum distance (denoted as u and v) between them in constant time. The main task of this procedure is to determine whether the two cells are resolvable or not. We call a pair of cells *resolvable* if both u and v fall into the same SDH bucket i. In the case when the two cells resolve into bucket i, we increment the distance count of that bucket by $n_A n_B$, where n_A and n_B are the number of particles in cell A and B, respectively. In the case of non-resolvable cells, we take one of the following actions:

- (1) Go to the next density map with higher resolution and resolve all children of A with those of B, or
- (2) If it is the leaf-level density map, compute every distance between particles of A and B and update the histogram accordingly.

To get the complete SDH of the MS system, the algorithm executes the RESOLVETWOCELLS procedure for all pairs of cells on a given density map DM_k (the DM where the diagonal of a cell is smaller than or equal to the bucket width w). So, basically, the algorithm calls RESOLVETWOCELLS recursively (i.e., action (1) above) till it reaches the leaf level of the tree (i.e., action (2) above). Via a geometric modeling approach, we have proved that the time complexity of DM-SDH is $O(N^{\frac{2d-1}{d}})$ [9], where d is the number of dimensions. But the more exciting news is that DM-SDH can be enhanced to approximately compute SDH with a time complexity of $I(\frac{1}{\epsilon})^{2d-1}$, where ϵ is an error bound and I is the number of pairs of cells in DM_k .⁴ This algorithm is named ADM-SDH.

The idea behind the ADM-SDH algorithm is to recursively call RESOLVETWOCELLS only for a predetermined number (m) of levels in the tree. If after visiting the m levels, there are unresolved pairs of cells,

 $^{^3\,}$ In this paper, we focus on 2D data to elaborate and illustrate the proposed ideas. The extension of our work to 3D space would be straightforward.

 $^{^4}$ There is also a $O(N\log N)$ cost for building the Quadtree.

heuristics is being used to greedily distribute distances into relevant SDH buckets. We will study the heuristics for distance distribution in Section 4. The beauty of this algorithm is: given a user specified error bound ϵ , our analytical model can tell what value of m to choose [42]. Although ADM-SDH is fast in regards to the data size N, its running time is very sensitive to the bucket width w. The main reason for this is: when wdecreases by half, we have to start the algorithm from the next level of the tree. As a result, the number of cells in the starting level DM_k increases by a factor of 2^d and I increases by a factor of 2^{2d} . Since the SDH is a discrete approximation of a continuous distribution of the distances in the MS system, more information is lost with the increase of w. In practice, scientists prefer smaller values of w so that there are a few hundred buckets in the SDH. In this paper, we present an efficient and accurate multi-frame SDH computing algorithm whose performance is insensitive to **both** N and w. This new algorithm uses the same region quad-tree for data organization as in the DM-SDH and ADM-SDH algorithms.

4 SDH Computation Based on Spatial Uniformity

4.1 Algorithm Design

The DM-based algorithms depend heavily on resolving cells to achieve the desired accuracy. Only when we finish visiting m levels of the tree or reach the leaf nodes do we use heuristics to distribute the distances into relevant buckets. That is the main reason for the long running time. Our idea to remedy that problem is to greedily distribute distances between very large regions of the simulation space, even when no pairs of such regions are resolvable. In other words, we use heuristics for distance distribution as early as possible. However, the distribution of distances between two large regions may yield arbitrarily large errors. Therefore, the key challenge is to design a heuristic with high accuracy even under large regions.

Our first idea to address the aforementioned challenge is to take advantage of the spatial distribution of data points in the cells. As illustrated in Fig. 3: two cells have a distance range [u, v] that overlaps with three SDH buckets (i.e., from bucket *i* to *i*+2). A critical observation here is: if we knew the probability distribution function (PDF) of the point-to-point distances between cells A and B, we can effectively distribute the actual number of distances $n_A n_B$ into the three overlapping SDH buckets. Specifically, the total number of $n_A n_B$ distances will be assigned to the buckets based on the



Fig. 3 Distance range of non-resolvable cells overlaps with more than one bucket of the SDH $\,$

probability of a distance falling into each bucket according to the PDF. For the case in Fig. 3, the relevant SDH buckets and the number of distances assigned to them are as follows:

$$H[i], \qquad n_A n_B \int_u^{iw} g(t) dt \tag{1}$$

$$H[i+1], \quad n_A n_B \int_{iw}^{(i+1)w} g(t)dt \tag{2}$$

$$H[i+2], \quad n_A n_B \int_{(i+1)w}^{t} g(t)dt \tag{3}$$

where g is the PDF. The biggest advantage of the above approach is that the errors generated in each distance count assignment operation can be very low, and the errors will not be affected by the bucket width w, as long as the PDF is an accurate description of the underlying distance distribution [8]. Therefore, the main task of the proposed approach is to derive the PDF.

4.1.1 Methods for deriving the PDF

Note that in the work presented in [42], the distances are proportionally distributed into the three buckets based on the overlaps between range [u, v] and the individual buckets. Such a primitive heuristic, which is named PROP (short for "proportional"), implicitly assumes that the distance distribution is uniform within [u, v]. However, our experiments show that a typical distance distribution in MS data is far from being uniform. Hence, our proposed solution will naturally introduce less errors than the PROP heuristics adopted by ADM-SDH.

In general, the PDF of interest can be obtained by the spatial distribution of particles in the two relevant cells. The coordinates of any two particles - one from A and the other from B - can be modeled as two random vectors \mathbf{v}_A and \mathbf{v}_B , respectively. The distance between these two particles can also be modeled as a random variable D, and we have

$$D = ||\mathbf{v}_A - \mathbf{v}_B||. \tag{4}$$

Given that, if we know the PDFs of both \mathbf{v}_A and \mathbf{v}_B , the PDF of D can be derived via either one of the following two strategies:

- (1) generation of a closed-form via analyzing the PDFs of \mathbf{v}_A and \mathbf{v}_B as well as Eq. 4; or
- (2) Monte Carlo simulations using the PDFs of \mathbf{v}_A and \mathbf{v}_B as data generation functions.

In practice, it is difficult to get a closed-form PDF for D even when the particle spatial distributions follow a simple form. In Section 4.3, we present the results of our efforts in obtaining such a closed-form PDF for Dunder uniformly distributed \mathbf{v}_A and \mathbf{v}_B .

Monte Carlo simulations can help us obtain a discrete form of the PDF of the distance distribution, given the PDFs of the particle spatial distributions [31]. One important note here is that the method works no matter what forms the spatial distributions follow. However, to generate the particle spatial distributions, it is infeasible to test the MS dataset for all possible data distributions. Instead, we focus on testing if the data follows the most popular distribution in MS - spatial uniform distribution.⁵ As discussed in Section 1.2, natural systems often contain localized regions with uniformly distributed components.

4.1.2 Algorithmic details

Given the discussions, our proposed algorithm contains the following steps:

- (1) Identifying **uniform regions** in which particles are uniformly distributed;
- (2) Deriving the distance distribution PDFs between all pairs of uniform regions by:
 - Mathematical analysis towards a closed-form, or
 Monte Carlo simulations;
- (3) Assigning the actual distance counts in such regions following Eqs. 1–3.

One technical detail skipped is that we also need to assign intra-cell distances to the first few buckets of the SDH, as step(2) only handles inter-cell distances. In particular, given a cell A with diagonal length of q, the distances between any two particles in A fall into the range [0, q], and can be modeled as the following random variable:

$$D' = ||\mathbf{v}_A - \mathbf{v'}_A|| \tag{5}$$

where $\mathbf{v'}_A$ is an independent and identically distributed variable to \mathbf{v}_A . Let us further assume the range [0, q]



Fig. 4 Sub-trees of nodes P and Q with their leaf nodes

overlaps with buckets 0 to j. Then we can follow the same idea shown in Eqs. 1-3 to assign the distance counts of cell A into the relevant buckets:

$$H[0], \quad \frac{n_A(n_A-1)}{2} \int_0^w g_{D'}(t)dt \tag{6}$$

$$H[1], \quad \frac{n_A(n_A - 1)}{2} \int_w^{2w} g_{D'}(t) dt \tag{7}$$

$$H[j], \quad \frac{n_A(n_A-1)}{2} \int_{(j-1)w}^q g_{D'}(t)dt \tag{8}$$

where $g_{D'}(t)$ is the PDF for random variable D', and can also be generated by mathematical analysis or approximated by Monte Carlo simulations.

Finally, to complete the above algorithm, all pairs of cells containing at least one non-uniform cell, will be processed using the PROP heuristic – same as in ADM-SDH. Physical study of molecular systems have shown that it is normal to see a small number of large uniform regions covering most of the particles, leaving only a small fraction of particles in non-uniform regions [3,5]. This is also verified by our experiments using real MS datasets (Section 9). This translates into high efficiency of the proposed algorithm. Furthermore, the time complexity of steps (1) and (2) is unrelated to the bucket size w. The time needed for step (3) is linearly and negatively related to w.

In the remainder of this section, we present technical details in implementing the different steps of our algorithm.

⁵ Particle spatial distribution is different from the distribution of distances between particles.

4.2 Identification of Uniform Regions

The first problem related to this topic is: given a spatial region (represented as a quad-tree node), how do we test if it is a uniform region? We take advantage of the chi-square (χ^2) goodness-of-fit test to solve this problem. Here we show how the χ^2 test is formulated and implemented in our model. A brief introduction to this statistical tool and justification of its applicability to our problem can be found in Appendix A.

Definition 1 Given a cell Q (i.e., a tree node) in the DM-tree, we say Q is uniform if its probability value p in the chi-square goodness-of-fit test against uniform distribution is greater than a predefined bound α .

To obtain the *p*-value of a cell, we first need to compute two values: the χ^2 value and the degree of freedom (df)of that particular cell. Suppose cell Q resides in level k of the DM-tree (see Fig. 4). We go down the DMtree from Q till we reach the leaf level, and define each leaf-level descendant of Q as a separate **category**. The intuition behind the test here is: Q is uniform if each category contains roughly the same number of particles. The number of such leaf-level descendants of cell Q is 4^{t-k} , where t is the leaf level number. Therefore, the df becomes $4^{t-k} - 1$. The observed value, O_j , of a category j is the actual particle count in that leaf cell. The expected value, E_j , of a category is computed as follows:

$$E_j = \frac{Total \ Particle \ Count \ in \ Cell \ Q}{\# \ of \ leaf \ level \ descendants \ of \ Q} = \frac{n_Q}{4^{t-k}} \quad (9)$$

Having computed the observed and expected values of all categories related to Q, we obtain the χ^2 test score of cell Q through the following equation:

$$\chi^2 = \sum_{j=1}^{4^{t-k}} \frac{(O_j - E_j)^2}{E_j} \tag{10}$$

Next, we feed these two values, the χ^2 and the df, to the R statistical library [41], which computes the pvalue. We then compare the p-value to a predefined probability bound α (e.g., 0.05). If $p > \alpha$, we mark the cell Q as uniform, otherwise we mark it as non-uniform. Note that the χ^2 test performs poorly when the particle counts in the cells drop bellow 5 [20]. But, we already had similar constraint in our algorithm while building the DM-tree, essentially making the cells in the leaf level contain more than 4 particles. Hence, we choose leaf level nodes as the categories in the test.

To find all the uniform regions, we traverse the DMtree starting from the root and perform the above χ^2 test for each node we visit. However, once a node is **Algorithm** MARKTREE(node Q, level a)

```
0 checkUniform(Q, a)
```

```
1 if Q is NOT uniform
```

```
2 then for each child B_i of cell Q: i := 1...4
3 MarkTree (B_i, a + 1)
```

Procedure CHECKUNIFORM(node Q, level a)

- $\begin{array}{ll} 0 & \mbox{Go to leftmost leaf level } (t) \mbox{ descendent of } Q \\ 1 & \mbox{ for } k = 1 \mbox{ to } 4^{t-a} \end{array}$
- 2 $\chi^2 := \chi^2 + \frac{(O_k E_k)^2}{E_k}$
- 3 Get $pval(\chi^2)$ using $\stackrel{L_k}{R}$ library
- 4 **if** pval > significance value α
- 5 **then** mark Q as uniform
- $6 \quad {\bf else} \ {\rm mark} \ Q \ {\rm as \ not \ uniform}$

Fig. 5 Marking uniform regions

marked uniform, there is no need to visit its subtree. The pseudo code shown in Fig. 5 represents this idea – to find all uniform regions, we only need to call procedure MARKTREE with the root node of the DM-tree as input.

4.3 Analysis of the PDF

In practice, it is difficult to get a closed-form PDF for D even when the particle spatial distributions follow a simple form. There has been some work done in [13] that addresses one special cases: tackling the distribution of distance between points within a unit square – this can be seen as a case of variable D' shown in Eq. (5). The distribution of random variable D is also studied in [2] under the special case that \mathbf{v}_A and \mathbf{v}_B are from two *adjacent* unit squares. Both work show closed-form and such formulae can be used in our algorithm. Since the formulae are complex, we list their results in Appendix B. To the best of our knowledge, there has not been any work that achieved derivation of a closed-form for the general cases.

In this part, we show the results of our efforts in obtaining an approximate closed-form for the general case: finding distance distribution between points in any two cells. The main claim is: if the data points in cell A and cell B are uniformly distributed, then the square of the distance between the two cells' points can be approximated by a **Noncentral chi-square** distribution and the distribution is not related to the number of points in cell A or B.

To shed more light on the claim, we take a look at two randomly chosen cells A and B of same size (i.e., from the same level of the DM tree). We start by assuming that the particles in the cells are uniformly

distributed. Our goal here is to give a representation of the PDF of the distance between points in such two cells. Let us choose two random points P_A and P_B , from cell A and B, and denote the coordinates of P_A and P_B as (X_{PA}, Y_{PA}) and (X_{PB}, Y_{PB}) , respectively. The square of the distance D between these two points can be expressed with the following equation:

$$D^2 = |X_{PA} - X_{PB}|^2 + |Y_{PA} - Y_{PB}|^2.$$

Since the points are chosen randomly, their coordinates can be regarded as random variables. Furthermore, $|X_{PA} - X_{PB}|$ and $|Y_{PA} - Y_{PB}|$ can be viewed as random variables that follow a triangular distribution. But using triangular distribution would make the result and the analysis really hard (if not impossible) to achieve. In order to ease the analysis process we will approximate the triangular distribution with a normal distribution. So, naturally, we continue by first figuring out how much error will be introduced by such approximation. The following subsection shows that the introduced error is only 10%.

4.3.1 Approximating Triangular with Normal distribution

Lemma 1 If X and Y are independent random variables uniformly distributed on (a,b) and (c,c+b-a), and $c \ge a$, then Y - X is a triangular random variable and can be regarded as a normal random variable with total variation distance 0.1.

Proof The probability density of X is

$$f(x) = \frac{1}{b-a}, \quad a < x < b$$
 (11)

and the probability density of Y is

$$g(y) = \frac{1}{b-a}, \quad c < y < c+b-a$$
 (12)

There are two cases to be considered: (1) when c is equal to a; and (2) when c is greater than a.

Case 1 (c = a): The probability density of Y - Xcan be calculated as follows

$$f_{Y-X}(z) = \int_{c}^{c+b-a} f(y-z)g(y)dy$$
 (13)

When 0 > z > a-b, the probability density of Y-X can be computed as follows

$$f_{Y-X}(z) = \frac{b-a+z}{(b-a)^2}$$
(14)



Fig. 6 The distribution of Y-X

When 0 < z < b-a, the probability density of Y-X can be computed as follows

$$f_{Y-X}(z) = \frac{b-a-z}{(b-a)^2}$$
(15)

Case 2 (c > a):

When c - b < z < c - a, the probability density of Y - X can be computed as follows

$$f_{Y-X}(z) = \frac{b-c+z}{(b-a)^2}$$
(16)

When c - a < z < c + b - a, the probability density of Y - X can be computed as follows

$$f_{Y-X}(z) = \frac{c+b-2a-z}{(b-a)^2}$$
(17)

The probability density of Y - X is illustrated in Figure 6.

Now, let us take a look at a different random variable Q. Assuming Q is a normal random variable with parameters $(c-a, \frac{(b-a)^2}{6})$, the probability density of Q can be written as follows:

$$f_Q(x) = \frac{\sqrt{6}e^{\frac{-6(x-c+a)^2}{2(b-a)^2}}}{\sqrt{2\pi}(b-a)}$$
(18)

Let $u = \frac{x - (c - a)}{b - a}$, then the probability density of Q can be rewritten as follows

$$f_Q(u) = \frac{\sqrt{6}e^{-3u^2}}{\sqrt{2\pi}}$$
(19)

Let $v = \frac{z - (c - a)}{b - a}$. Then, the probability density of Y - X can be rewritten as follows

$$f_{Y-X}(v) = Min\{1+v, 1-v\}, \quad -1 < v < 1$$
(20)

Now let us study how well the normal distribution approximates the triangular distribution. Let Pbe the triangular distribution with PDF p(x) given by Eq. (20) and Q is the normal distribution $N(0, \frac{1}{6})$, the PDF of which is $q(x) = \frac{\sqrt{6} \exp\{-3x^2\}}{\sqrt{2\pi}}, x \in (-\infty, +\infty).$



Fig. 7 The difference between normal and triangular distributions

Note that the first two moments of the two distributions are exactly the same: E[P] = E[Q] = 0, and $\operatorname{var}[P] = \operatorname{var}[Q] = \frac{1}{6}$. Although these indicate the similarity between the two distribution, we still want to quantify how close the two probability measures are.

A natural measure of the difference between the two probability measures P and Q is the **total variation distance** defined as:

$$V(P,Q) \triangleq \sup_{A \in \mathscr{F}} |P(A) - P(A)| = \frac{1}{2} \int_{R} |p(x) - q(x)| dx,$$

where \mathscr{F} is the σ -field upon which the probability space is defined. Note that P is absolute continuous with respect to Q in our case, then we have

$$V(P,Q) = \frac{1}{2} \int_{R} \left| \frac{p(x)}{q(x)} - 1 \right| q(x) dx$$

Numerical computation shows that V(P,Q) = 0.1012in this case.

Following the above reasoning, we conclude that we can use normal distribution instead of triangular distribution, introducing an error of about 10%.

Using the aforementioned findings, we now regard the differences $|X_{PA} - X_{PB}|$ and $|Y_{PA} - Y_{PB}|$ as random variables with normal distribution. In the following subsection, we continue with the proof of our main claim. Here we show that the square of the distance can be viewed as random variable with non-central chisquared distribution.

4.3.2 Distance distribution of particles in two cells A and B

As we know, if X_{PA} and X_{PB} are independent random variables uniformly distributed on (a, b) and (c, c+b-a)respectively, $(c \ge a)$ then $X_{PB} - X_{PA}$ follows a triangular distribution that we saw can be approximated with normal, introducing an error of not more than 10%. Knowing this, $X_{PB} - X_{PA}$ can be regarded as a normal random variable with parameters $(c-a, (b-a)^2/6)$. Similarly, $Y_{PB} - Y_{PA}$ can be regarded as a normal random variable with parameters $(c'-a', (b-a)^2/6)$. Since $(b-a)^2/6$ is a constant, which is noted as σ^2 , we can write the following equation for the distance D between the two points P_A and P_B :

$$\frac{D^2}{\sigma^2} = \frac{(X_{PB} - X_{PA})^2}{\sigma^2} + \frac{(Y_{PB} - Y_{PA})^2}{\sigma^2}$$
(21)

As it is known, the right hand side of the above equation is a Noncentral chi-square distribution. This means that the distances between the two cells' points can be described as a Noncentral chi-square distribution with the parameters $(2, \lambda)$, where λ can be defined as follows:

$$\lambda = \frac{(c-a)^2}{\sigma^2} + \frac{(c'-a')^2}{\sigma^2}$$
(22)

where c - a and c' - a' are the means of the two normal distributions.

Note that, since our discussions started with the only assumption that points in A and B are uniformly distributed, the parameters of above PDF have no relationship with the actual number of points in cell A or cell B.

So, our conclusion is that the square of the distance between any two points from two cells follows (can be approximated to) a Noncentral chi-square distribution. Since the PDF of a Noncentral chi-square distribution has a closed form [12], the PDF of D (i.e., the square root of the Noncentral chi-square) can be obtained through Jacobian transformation. However, we stop here after obtaining the (approximated) PDF of D^2 since it can already be used to guide distance distributions in our algorithm with minor tweaks. Recall the scenario in Figure 3: the share of distance counts that should go into bucket *i* is now $\int_{u^2}^{i^2w^2} h(t)dt$ where h(t) is the PDF of the Noncentral chi-square. The other buckets can be treated in a similar way.

It is our belief, based on the thorough work we have done on this matter, that to get an explicit and more accurate closed form for the distribution of the distances between the points of the cells is a really challenging, if not impossible to solve, problem.

4.4 Monte Carlo Simulations

The distribution of distances between a pair of cells, say A and B, can be determined based on their spatial distribution of particles, by running Monte Carlo simulations. Monte Carlo simulation is a way to model a phenomenon that has inherent uncertainty [31]. If the spatial distributions of particles in A and B are known to be uniform, the simulations can be done by sampling (say n_s) points independently at random from uniform distributions within the spatial ranges of A and B. Then, the distance distribution is computed from the points sampled in both cells. A *temporary* distance histogram can be built for this purpose. All n_s^2 distances are computed (brute-force method), and put into buckets of the temporary histogram (e.g., those overlapping with [u, v] in Fig. 3) accordingly. The final proportions of each bucket in the temporary histogram will fulfill our needs in step (3) of the algorithm.

Sufficient number of points are needed to get reasonably high accuracy of the SDH generated [8]. The cost of running such simulations can be high if we were to perform one simulation for each pair of uniform regions. This, fortunately, is not the case. First, let us emphasize that the simulations are not related to the number of particles (e.g., n_A and n_B) in the cells of interest - the purpose is to approximate the PDF of distance distribution. Second, and most importantly, the same simulation can be used for multiple pairs of cells in the same density map, as long as the two cells in such pairs have the same relative position in space. A simple example is shown in Fig. 2: cell pairs (A, B) and (A, B_1) will map to the same range [u, v] and can definitely use the same PDF. A systematic analysis of such sharing is presented in following theorem.

Theorem 1 The number of distinct Monte Carlo simulations performed for all pairs of cells in a density map of M cells, is O(M).

Theorem 1 says that, for the possible $O(M^2)$ pairs of uniform regions on a density map, there are only a linear number of simulations that need to be run. Furthermore, as we will see later (Section 5), the same cells exist in all the frames of the dataset, therefore, a simulation run for one frame can be shared among all frames. Given the above facts, we can create a lookup table (e.g., hash-based) to store the simulation results to be shared among different operations when a PDF is required.

Remark 1 If we were given the PDF of the random variable D and use the integration of the PDF to guide distance distribution in step (2) of our algorithm, the number of distinct integrations is also O(M).

5 SDH Computation Based on Temporal Locality

Another inherent property of the MS is that the particles often exhibit temporal locality, and such temporal

property can be utilized to compute the SDH of consecutive frames even faster. The existence of temporal locality is mainly due to the physical properties of the particles in most of the simulation systems. More specifically, such properties can be observed at the following two levels:

- (1) Particles often interact with each other in groups and move randomly in a very small subregion of the system;
- (2) With particles moving in and out of a cell, the total number of particles in that cell does not change much over time.

5.1 Basic Algorithm Design

We discuss the algorithm in terms of only two frames f_0 and f_1 , although the idea can be extended to an arbitrary number of frames. Suppose DM-trees T_0 and T_1 are built for the two frames f_0 and f_1 , respectively. Since the DM-trees are built independently from the data they hold, the number of levels and cells, as well as the dimensions of corresponding cells in both DM-trees will be the same. First, an existing algorithm (e.g., DM-SDH or ADM-SDH) is used to compute the SDH H_0 for the base frame f_0 . Then we copy the SDH of frame f_0 to that of f_1 , i.e., $H_1 = H_0$. The idea is to modify the initial value of H_1 to reach its correct form by only processing cells that do not show temporal locality.

Let DM_k^0 and DM_k^1 be the density maps, at level k, in their respective DM-trees T_0 and T_1 . We augment each cell in DM_k^1 with the ratio of particle count of that cell in DM_k^1 to the particle count of the same cell in DM_k^0 . A density map that has such ratios is called a ratio density map (RDM). The next step is to update the histogram H_1 according to the ratios in the RDM. Let r_A and r_B ($A \neq B$) be the density ratios of any two cells A and B in the RDM, we have two scenarios:

Case 1: $r_A \times r_B = 1$. In this case, we do not make any changes to H_1 . It indicates that the two cells A and B contributed the same (or similar) distance counts to the corresponding buckets in both histograms H_0 and H_1 .

Case 2: $r_A \times r_B \neq 1$, which indicates that some changes have to be made to H_1 . Specifically, we follow the PROP heuristic, as in ADM-SDH, to proportionally update the buckets that overlap with the distance range [u, v]. For example, as shown in Fig. 3, consider the distance range [u, v] overlapping three buckets i, i + 1, and i + 2. The buckets and their corresponding count



Fig. 8 Grouping cells with equal density ratios by sorting the cell ratios in the RDM

updates are given in Eqs. 23–25.

$$H_1[i], \quad \left(n_A^1 n_B^1 - n_A^0 n_B^0\right) \frac{iw - u}{v - u} \tag{23}$$

$$H_1[i+1], \quad \left(n_A^1 n_B^1 - n_A^0 n_B^0\right) \frac{w}{v-u} \tag{24}$$

$$H_1[i+2], \quad \left(n_A^1 n_B^1 - n_A^0 n_B^0\right) \frac{v - (i+1)w}{v - u} \tag{25}$$

where n_A^0 and n_B^0 are counts of particles in cells A and B, respectively, in density map DM_k^0 of frame f_0 . Similarly, n_A^1 and n_B^1 are counts of particles in corresponding cells of density map DM_k^1 in frame f_1 . Note that we have $n_A^1 = r_A \cdot n_A^0$ and $n_B^1 = r_B \cdot n_B^0$. The total number of distances to be updated in the buckets is $n_A^1 \times n_B^1 - n_A^0 \times n_B^0$. This actually gives us the number of distances changed between cells A and B of density map DM_k , going from frame f_0 to frame f_1 .

5.2 Algorithmic Details

An efficient implementation of the above idea requires all pairs of cells that satisfy the **Case 1** condition to be skipped. In other words, our algorithm should **only** process the Case 2 pairs, without even checking whether the product of two cells is 1.0 (explained later). The histogram updates can be made efficiently if cells with equal or similar density ratios are grouped together. Our idea here is to store all the ratios in the RDM in a sorted array (Fig. 8). The advantage in sorting is that the sorted list can be used to efficiently find all pairs of cells with ratio product of 1.0. In other words, for any cell D with density ratio r_D , find the first cell E and the last cell F in the sorted list with ratios $1/r_D$, using binary search. Then, pair cell D with all other cells except the cells between E and F in the sorted list. Fig. 8 shows an example of a cell (D_1) with ratio 1.0 – we mark the first cell E_1 and the last cell F_1 with ratio of 1.0. Then we pair D_1 with rest of the cells in the list. Take another example of cell (D_2) with ratio 0.2: we will effectively skip all the cells (E_2 to F_2) with ratio 5.0 (as 1/0.2 = 5.0), and start pairing D_2 with those cells that do not have ratio 5.0 (to the left of E_2 and right of F_2).

In practice, a tolerance factor ϵ can be introduced to the **Case 1** condition such that the cells with ratio product within the range of $1.0 \pm \epsilon$ are skipped from the computations. While saving more time by allowing more cell pairs untouched, the factor ϵ can also introduce extra errors. However, our analysis in Section 7 shows that such errors are negligible. Our experimental results (Fig. 14(b)) show that there are a large number of pairs of cells whose density ratio products are around 1.0, thus providing sufficient savings of computation.

Same as in Section 4.1.2, there are also intra-cell distances to be processed. Again, we assume the cell of interest has diagonal length q, and the distance range [0,q] overlaps with buckets $0, 1, \ldots, j$. If an individual cell is with an RDM of 1.0, nothing needs to be done. For those cells whose RDM is not 1.0, the following rules are used to update the counts.

$$H_1[0], \quad \left[\frac{n_A^1(n_A^1-1)}{2} - \frac{n_A^0(n_A^0-1)}{2}\right]\frac{w}{q} \tag{26}$$

$$H_1[j-1], \quad \left[\frac{n_A^1(n_A^1-1)}{2} - \frac{n_A^0(n_A^0-1)}{2}\right]\frac{w}{q} \tag{27}$$

$$H_1[j], \quad \left[\frac{n_A^1(n_A^1-1)}{2} - \frac{n_A^0(n_A^0-1)}{2}\right] \frac{(j-1)w}{q} (28)$$

6 Putting Both Ideas Together

The continuous histogram processing is sped up by utilizing both spatial uniformity and temporal locality properties. An overview of the technique is shown in the flow diagram of Fig. 9. The left branch (decision $A \equiv B$) is to compute the intra-cell distances. In the right branch we check the locality property of every pair of cells before checking for uniform distribution of the particles. Any pair that satisfies the locality property is skipped from further computations. The pairs that fail the locality property check are tested for the uniformity property. Based on the results of the check, subsequent steps are taken and the histogram buckets are updated.

The Monte Carlo simulation step introduced in our algorithm is expensive when computing SDH of a sequence of frames. As mentioned in Section 4, the cost can actually spread over when we are processing a sequence of frames. It is an interesting fact that the tree building process is such that a cell in the DMs of same



Fig. 9 Steps in dealing with two cells of the composite algorithm for computing SDH

level in all frames is of same dimensions. Therefore, a simulation done once can be reused in all other frames. Given a pair of cells A and B and their respective distance range [u, v], we compute the proportions of distances that map to each bucket covered by [u, v] through Monte Carlo simulation. For each distinct [u, v] range, we store such (and only such) proportions of distance distributions in a universal hash table.

For every pair of uniform cells that do not resolve and have distance range [u, v], we look into the hash table to get the proportions to distribute the distances into buckets. If an entry is available in the hash table, we use it directly. Otherwise, a new simulation is done and proportions are calculated. This new simulation information is stored in the hash table. The hash table is universal and is used for computing the histogram of all the frames for a given bucket width.

The same strategy can be followed if we were to use closed-form PDFs (if available) to determine the proportions of distances.

To simplify the implementation, one decision we made was to choose a level k in the DM-tree and process cells on that level only (instead of working on uniform regions on different levels). We need a level that balances both SDH computation time and the error – choosing a level close to the leaves may increase

the time, while a level close to the root will introduce higher errors in the SDH. An important feature of our algorithm is that the user can choose a level to run the algorithm according to her tolerance of the errors. Such choices can be made beforehand by analysis as discussed in Section 7.1.3. Note that all the cells in the DM-tree that are uniform are marked before the continuous SDH processing begins.

The proposed technique is completely based on the general temporal and spatial uniformity of the data set. Such cell-wise uniformity is not only observed in MS, but also in many traditional spatiotemporal database applications [39]. Hence, it can be applied to very different data sets such as crowd of people and stars in astronomical studies.

7 Performance Analysis

The performance (running time and errors) of the proposed technique in utilizing the spatio-temporal property of the data is analyzed in this section.

7.1 Analysis About Spatial Uniformity

7.1.1 Time analysis

The running time of the algorithm utilizing only the spatial uniformity property is contributed by the following factors:

- (1) Quad tree construction time $O(N \log N)$ where N is the number of particles in simulation;
- (2) Identification of uniform regions. This can also be bounded by $O(N \log N)$, as the count in each leaf node is used for at most $\log N$ chi-square tests;
- (3) Distribution of distances into buckets; For this, all pairs of cells on a DM need to be computed in a DM with M cells, the time is $O(M^2)$.
- (4) Monte-Carlo simulations that require $O(MT_s)$ time according to Theorem 1. Here T_s is the time of each individual simulation.

Theoretically, the first two costs will dominate as their complexity is related to system size N. In practice, the $O(M^2)$ time for factor (3) can dwarf others if we choose a density map on the lower levels of the quad tree - M approaches N when the level gets lower. Recall that this will happen to the ADM-SDH algorithm when the bucket width w gets smaller. Our following analysis and experiments will show that M is under control in our algorithm.

Factor (4) is also worth a special note. Although the simulation time T_s can be regarded as a constant (as it

is unrelated to both N and w), a larger number of points in the simulation is preferred for better accuracy. Thus, it is essential to study how many data points we have to simulate in order to reach desired accuracy. Such analysis is shown in Section 7.1.2.

7.1.2 Error analysis

Based on the sources, two types of errors are introduced by utilizing the spatial uniformity feature:

- I. error (e_u) by pairs of cells that are both uniform, and
- II. error (e_a) by those with at lease one non-uniform cell.

Type I error is basically the simulation error, i.e., the expected percentage of distances put into the wrong buckets when both cells have uniformly distributed data points. According to the Law of Iterated Logarithm (LIL) [7], such error is up to the order of $\left(\frac{S_m}{\log \log S_m}\right)^{-1/2}$, where S_m is simulation size. Since we compute the Euclidean distance between two randomly selected points which are uniformly distributed in the two cells, we have $S_m = n_s^2$, where n_s is the number of points simulated in each cell. Clearly, the error drops dramatically with the increases of n_s . Considering a scenario where n_A and n_B are of the order of 10^2 , the simulation error is slightly smaller than the order of 10^{-2} . In other words, we can effectively control the Type I error without suffering from a heavy simulation overhead.

The Type II error is obviously no greater than the error achieved by the PROP heuristic. It is hard to get a tight error bound when the distribution of points in a cell is not uniform. But it is easy to see that the error for one single distribution using PROP can be arbitrarily large. Unlike the Type I error, error in this category cannot be controlled. At this point, we can at least conclude that, due to the small Type I error, our algorithm will be more accurate than existing solutions based on PROP, such as ADM-SDH [42].

An important note here is that our analysis has so far concentrated on the errors introduced in an individual distribution operation (i.e., between one pair of cells). However, our work [21] has revealed the fact that errors generated by different pairs of cells can cancel out, and reduce the error in the whole SDH to a great extent. We call such a phenomenon *error compensation*. In particular, our qualitative study shows that the error (at the entire SDH level) caused by PROP can be loosely bounded by 10%. Since this is not a tight bound, we expect to see much smaller errors in practice, as shown in our experimental results for the ADM-SDH algorithm (Section 9.2). For the same reason, the effects of Type I error can also be reduced by error compensation, making the Type I error a negligible quantity.

7.1.3 Error/performance tradeoff

Given the above analysis, we show our algorithm is *tunable* in that the user can choose a level of DM-tree to get a desired error guarantee. Suppose p_u is the fraction of pairs of cells that are uniform on a given level, the total error ξ produced by our algorithm based on spatial uniformity is

$$\xi \le e_u p_u + e_a (1 - p_u) \tag{29}$$

A remark here is: as compared to ADM-SDH that is based on PROP heuristics, our algorithm shows an advantage in accuracy: error will be lower by $(e_a - e_u)p_u$.

From the above equation, we can also solve p_u to obtain a guideline on the level of the DM tree from which we run the algorithm:

$$p_u \ge \frac{e_a - \xi}{e_a - e_u} \tag{30}$$

In other words, a user will choose to work on a DM where the fraction of uniform cells is at least $\sqrt{p_u}$, in order to get an error lower than ξ .

One special note about p_u is: defined as the fraction of **actual** uniform cell pairs, p_{μ} is smaller than the percentage of cell pairs marked as uniform by our algorithm shown in Fig. 5. This is because it is not a deterministic decision to mark a cell uniform, and cases of false positive can happen. In marking the cells, the chance of getting a false positive consists of the approximation error of the Pearson's χ^2 test statistic [7] and the probability bound α used in the test. The test statistic error is up to the order of $O_t^{\frac{1-\nu}{\nu}}$, where ν is degree of freedom and O_t is the number of observations in χ^2 test. In our environment, O_t tends to be a large number, as we often see large uniform regions. The α value is user tunable and usually set around 5%. When ν is sufficiently large, the error in marking a cell uniform is $\gamma = \alpha + 1/O_t \approx \alpha$. Therefore, if the percentage of pairs of cells marked uniform by our algorithm is p'_u , we have

$$p_u = (1 - \gamma)^2 p'_u \approx (1 - \alpha)^2 p'_u$$

7.2 Analysis About Temporal Locality

7.2.1 Time analysis

The running time is determined by the number of cell pairs that do not satisfy the temporal locality condition, i.e., ratio products are not in the range of $1.0\pm\epsilon$. Due to the sorted list of ratios in the RDM, all cell pairs satisfying the above condition are skipped by the algorithm. Suppose p_r is the fraction of such cell pairs, only $(1-p_r)$ pairs of cells need to be processed by the algorithm. The sorting and searching of the cells can be performed in $O(M \log M)$ time. Hence, the running time of the algorithm is bound by $(1 - p_r)T + O(M \log M)$ where T is the time for processing the base frame. In other words, by utilizing the temporal locality, we achieve a $(1 - p_r)$ -factor improvement in running time.

7.2.2 Error analysis

We tackle this by studying the *extra errors* our algorithm generates for a frame f_1 on top of those in the base frame f_0 . The error introduced when utilizing the temporal locality can be categorized based on two cases:

- 1. temporal locality property is satisfied, and
- 2. temporal locality property is *not* satisfied.

Case 1: Error is produced by temporal locality property only when the cell pairs satisfy the condition $r_A \times r_B = 1.0 \pm \epsilon$. A small error equal to the fraction ϵ is introduced. When the fraction $\epsilon = 0$, there is no change in the number of distances between the two cells. In both situations, a negligible error, very hard to compute, is produced due to small change in position of the points. The fraction ϵ is negligible when the pairs of cells have uniformly distributed points in both the frames f_0 and f_1 . Actually, the small movement of particles has minimal effects on the distance distribution.

Case 2: This case will not cause any additional errors. When the temporal locality condition is not satisfied for a pair of cells in f_1 , we update the histograms as if we are running the algorithm for the base frame therefore the error will be on the same level as in the base frame. On the other hand, we do not save any processing time in such cases.

From the above analysis, we conclude that the error in the derived frame will be on the same level as that of the base frame. Note that this conclusion should be interpreted in a statistical way.

8 SDH Computation on Graphics Processors

In this section, we look at the basic architecture of the GPUs and their programming paradigms. Then we modify our algorithm of utilizing spatiotemporal uniformity to map onto the GPU programming environment. Our discussions, however, will focus on how to optimize our algorithm in a typical GPU architecture rather than a straightforward implementation. This is because the



Fig. 10 The basic architecture of modern graphics processors (GPUs)

GPU architecture is very different from that of CPUs thus code optimization requires special (and sometimes unintuitive) techniques. For example, the GPU hardware provides a hierarchy of programmable memories with heterogeneous capacity and performance. For that, the data can be organized, on these memories, in such a way that the access latency is minimized. We also study other possible optimizations to further improve the performance by using such memory. The complexity of the computations such as, instruction branching and memory access patterns, affect the advantages of such special hardware features.

8.1 GPU Architecture

The basic GPU architecture, for both NVIDIA [34] and AMD [4] products, is illustrated in Fig. 10. The GPU consists of many *multiprocessors* that execute instructions on a number of GPU cores in a SIMD (Single Instruction Multiple Data) manner at any given clock cycle. The GPU devices have a considerable amount of global memory with high bandwidth. For example, the NVIDIA GTX 570 we used has 15 multiprocessors, each of which encapsulates 32 GPU cores. It also has about 1.2 GB of global memory with a bandwidth of 152 GB/s.⁶ Apart from the global memory, the GPUs have programmable, very fast cache memory (called *shared memory*). This type of memory is on-chip and shared by all GPU cores in a single multi-processor. Since it is onchip the access latency is very low. In contrast to that, the global memory has high access latency (400 to 800 clock cycles [34]). Therefore, the access pattern should be optimized to reduce the overall latency caused by global memory.

 $^{^{6}\,}$ In high-end cards such as Tesla C2075, the amount of global memory can reach 6GB.

A large number of threads can be executed in SIMD fashion on the GPUs. The major difference between CPU and GPU threads is that the GPU threads have low creation and context-switch time. We follow the terminology of NVIDIA's compute unified device architecture (CUDA) [34] to describe the operation of GPU multiprocessors. A group of threads executing on a multiprocessor is called *block*. The blocks are scheduled dynamically on different multiprocessors. All threads within a block share all the resources, such as registers, L1 cache etc., available on the multiprocessor. An interesting feature of the memory in GPUs is that different threads in a block can read different memory locations simultaneously. This is achieved only when threads read consecutive memory locations. The underlying hardware groups the consecutive memory access requests into one access. This process is called *coalesced* access. How to fully utilize the available high bandwidth via coalesced access is another direction for code optimization.

8.2 Minimizing Branches in GPU Kernels

In the computation of SDH, different paths will be taken by the algorithm to process cells in a density map (Fig. 9). This causes a major problem in GPU implementation as conditional branches in the code can significantly degrade performance in SIMD architectures [43,35]. Therefore, the first code optimization strategy we take is to minimize the number of branches. The first branch can be easily removed: we process the cases of intracell computation ($A \equiv B$) and intercell computation separately. Note that the intercell computation is the one that dominates the running time of our algorithm (Section 7.1.1). Thus, we invest most of our efforts into optimizing such computations.

Processing pairs of cells: We follow the steps after the first conditional branch shown in the algorithm flow diagram (Fig. 9) to process the density map. The steps have to be followed for each pair of cells. By processing different pairs of cells in parallel, we can achieve improved performance. The cells of the chosen density map are placed in the memory such that consecutive threads can read consecutive cells through coalesced access mechanism. A pair of cells of the density map, to be processed, are assigned to a single thread of the GPU. Each thread processes distinct pair of cells. Each block (group of threads) executing on a multiprocessor processes different portions of the density map. Thus, parallel processing improves the performance of the algorithm.



Fig. 11 Grouping cells in global memory and loading into shared memory for improving performance

Precomputing Monte-Carlo simulations: When the distances between a pair of cells have to be distributed into different buckets, the algorithm has to make such decisions based on Monte Carlo simulations. This causes the large number of threads to diverge (at the "Entry found" condition box in Fig. 9), which could hinder the performance. These divergences can be eliminated if the simulations are run and their results are stored in the hash table beforehand. When processing pairs of cells, we send all requests to such information directly to the pre-built hash table. This is a feasible plan since Theorem 1 sates there are only a linear number of simulations to be run. By this, Monte-Carlo simulations would not hinder the performance. Section 8.4 explains details about efficient computation of simulations.

8.3 Memory Optimization and Issues

The speed of memory access can be improved by placing the cells of the density map in shared memory. Each thread can access distinct pairs of cells from the shared memory. One major obstacle in implementing this idea is the limited size of shared memory. Therefore, we pair cells by grouping them. Let M be the number of cells in a density map and shared memory can hold $2M_S$ cells. We divide the shared memory into two sections, each holding up to M_S cells. With M_S as the size for group of cells, we have $G_c = M/M_S$ number of groups out of M cells of the density map. Each CUDA block can process two groups of cells in shared memory. Fig. 11 shows the mechanism of processing these groups. First, the cells belonging to groups G_i and G_j are loaded into shared memory. One cell is chosen from each group and paired to process. This is repeated for all cells in G_i and G_i . Then, the cells within each group are paired and processed. Next, the second group G_j is evicted and a new group G_k is brought into the shared memory. This is repeated for all the groups of the density map until all the cells are processed. We can easily see that



Fig. 12 Illustrating bank conflicts in shared memory access on ${\rm GPU}$

such a cell grouping strategy can significantly reduce the number of global memory accesses.

There are some other issues with the utilization of shared memory for SDH computation:

- Bank conflicts: The shared memory is organized as banks in the hardware such that the threads read different banks in parallel. If threads read different addresses in the same bank, it gives rise to an access conflict called *bank conflict*. An example of bank conflicts is illustrated in Fig. 12.
- Computation intensive task: The operations of the algorithm, as shown in Fig. 9, are computation intensive rather than memory access intensive. Once, the information about cells is accessed, a large number of operations are performed. This shadows the latency involved in memory accesses.
- Thread divergence: Even after our efforts to reorganize the workflow in Fig. 9, there are still branches in the code. Once threads in the same block diverge, the hardware serializes the processing. But, the parallelism between blocks is maintained as they are assigned to different multiprocessors.

Due to the above issues, thread level parallelism may not be achieved in all cases. However, parallelism will be achieved at the multiprocessor level in the worst case.

8.4 Efficient Simulation

The cells of a density map have fixed dimensions. When a Monte-Carlo simulation is performed, we generate fixed number of random points in these cells and compute the distribution of distances between points of two cells. Such distribution is repeated for all pairs of cells for which distance distribution is required. However, according to Theorem 1, total number of actual simulations to be performed for a given density map is linear. Hence, the simulations can be performed on the GPU before the algorithm is executed. We utilize the shared memory of the multi-processors to optimize the simulations.

A set of random numbers are generated between range 0.0 to 1.0 in the shared memory for two cells. Instead of generating random numbers for every pair of cells, these are used for all the simulations required for processing SDH. Given two cells, of a density map, these random numbers are mapped to the boundaries of the cells. This operation does not require global device memory access, as the data is in very fast shared memory. Each block running on distinct multi-processor of the GPU can perform this mapping for different pairs of cells in parallel. The numbers are organized in the shared memory such that all the accesses belong to different banks.

Thus, the simulations can be performed efficiently by eliminating bank conflicts. Once we have the simulations performed, the distance distribution are computed and stored in a hash table. The hash table is stored in the global memory, due to limited size of the shared memory that is already occupied by the simulated points. The hash table is then used by the algorithm, eliminating the factors that would affect GPU performance in performing all need-based simulations.

9 Experimental Evaluations

9.1 Experimental Setup

The proposed continuous SDH computation algorithm was implemented in C++ programming language and tested on real MS data sets. The experiments were conducted on an Apple Xserve server with two Intel quadcore processors and 24 GB of physical memory. The Xserve was running OS X 10.6 Snow Leopard operating system. We tested the following algorithms to evaluate the performance of our approach.

- A1: The ADM-SDH algorithm presented in our previous work [42]. This method processes SDH frame by frame, and distributes the distances using PROP exclusively;
- A2: The algorithm utilizing only temporal locality to compute SDH continuously over multiple frames;
- A3: The algorithm utilizing only spatial uniformity to compute SDH frame by frame;
- A4: The algorithm utilizing both temporal locality and spatial uniformity to compute SDH continuously.

The running time of the algorithms on different data sets are measured for comparison, along with the errors introduced due to approximation. The errors are computed by comparing the approximate SDH results with the correct SDH of each frame. The error (in percentage) of each frame is calculated as

$$P_{error} = 100 \times \frac{\sum_{i=0}^{l-1} \left| H[i] - H'[i] \right|}{\sum_{i=0}^{l-1} H[i]}$$



Fig. 13 Percentage of the area of uniform regions at different levels of the DM tree

where H[i] and H'[i] are the correct and approximated distance counts in bucket *i* of the histogram, respectively.

Data Sets: Two datasets from different simulation systems were used for experiments. The first dataset consists of frames captured from a collagen fiber simulation system. The simulated system is made of 890,000 atoms and their positions are stored in a total number of 10,000 frames. The second dataset is collected from a simulation of cross membrane protein system with about 8,000,000 atoms and 10,000 frames. We randomly selected a chunk of 100 consecutive frames from the first dataset and 11 frames from the second dataset for our experiments. The main bottleneck in testing the algorithms is computing the correct histogram of the frames, needed to compute the error. Obtaining correct histogram is basically running the naive or DM-SDH algorithm, which is computationally expensive. Therefore, we could only get the correct histograms of 11 frames from the 8 million dataset (by naive approach in about 27 days!).

The percentage of cells with uniform data distribution (i.e., uniform regions) at different levels of the density map tree is shown in Fig. 13. The leaf level of the tree is not used to determine the uniformity, as very few particles fall into small cells. For both datasets, we started to see considerable amount of uniform regions at level 6 of the tree. Note that level 6 is still at the higher end of the tree (total number of levels is 9 for the smaller dataset and 11 for the larger one) and the total number of cells is only $4^6 = 4,096$. At level 8, the percentage of uniform regions is already over 90%. This confirmed the great potential of using spatial uniformity to save time in SDH processing.

We also observed significant temporal similarity among frames of both datasets. The success of utilizing the temporal similarity property depends on the total fraction of cells that exhibit such property. In fact, the



Fig. 14 Temporal similarity between two consecutive frames chosen randomly from the dataset of 890K atoms

running time of the technique is affected by the number of cell pairs (A, B) for which $r_A \times r_B = 1 \pm \epsilon$. Figs. 14(a) and 14(b) show the density of ratios and ratio products at each level of the DM-tree in two consecutive frames, chosen randomly from the dataset of 890,000 atoms. For all levels we tested, majority of the cells (cell pairs) show ratio (ratio product) that is close to 1.0. The number of cell pairs with ratio product of 1.0 increases as we descend down the tree.

9.2 Results of CPU Experiments

Main results: The average running time of all the algorithms for different bucket widths is shown in Fig. 15(a)and 15(c). It can be noted that the running time of A1 can be orders of magnitude longer than our proposed algorithms. The important observation to be made about algorithm A1 is that the running time increases dramatically with the decrease of w (note the logarithmic scale). Method A2 is similar to A1 but, utilizes temporal locality while working on only one level. When the bucket width is small, both methods work on lower tree levels, with small number of atoms in the cells. The utilization of locality gives scope to save some running time in A2. Unlike the first two methods, the time spent by methods A3 and A4 does not change much with the change of bucket width w. The data size however, limits the levels at which the algorithms work. Changing levels would affect the running time. The algorithms run at tree levels 6 and 7 for 890K and 8 million data set, respectively.⁷ Such levels are chosen to ensure that 80% of the area is covered by uniform regions (see Fig. 13). We generate 75 points from each of the two cells in Monte Carlo simulations - this number is chosen based

⁷ In [29], we run experiments on levels 5 and 6 of these two datasets, respectively, and very similar results are reported.



Fig. 15 Comparison of average running time and percentage errors of different algorithms. Both algorithms A3 and A4 process level 6 of the DM tree. (a)–(b) The results from 890,000 atom dataset. (c)–(d) Results from 8 million atom dataset

on our empirical results about sufficient simulation size (Fig. 19). Note that the average running time presented here have amortized all "start-up" costs including that for running Monte Carlo simulations and spatial uniformity test. The running time for larger bucket width is close to algorithm A1 and A2. This is because, in A1 and A2, the processing level is closer to root than the (fixed) level of tree used in algorithms A3 and A4. When we choose to have A3 and A4 run on a higher level, their time will clearly beat A1 and A2, as we have shown in [29]. The performance of A2 under smaller bucket width is bad because it works for lower levels of the tree, and the temporal locality is weak due to the small number of particles in each cell. For example, if there are 4 atoms in a cell in the base frame and one atom moves out of it in the derived frame, the density ratio is as low as 3/4 = 0.75.

The average errors (in percentage) of each method are shown in Fig. 15(b) and 15(d) for different values of w. The errors rendered by A3 and A4 are always lower than those by method A1. However, the errors of A2 are slightly higher than A1 for small bucket width. The number of distances to be distributed between two cells is very small, as the algorithm works close to leaf level. Therefore, by utilizing the temporal locality property the small errors are added on top of the PROP method applied for other cell pairs. Although method A4 is faster than A3, the price for that is an error rate that is slightly higher, as we expected based on our analytical results (Section 7.2). However, it provides a good tradeoff as the improvement of performance is of larger magnitude than the loss of accuracy. The method A3 stands clear winner in accuracy of the results. The distance distribution curve computed by Monte Carlo simulations diminishes the error that would have been introduced by heuristically distributing distances as in A1. The errors in method A1 stay low (still equal to

or higher than other methods) for smaller bucket width but goes higher under larger w values. The reason being, proportions for small buckets are almost similar in all the algorithms. Number of distances that are in the range of very small buckets are few and therefore their proportional distribution are not much different. Hence, the error is low. With the increase of bucket width, A1 would end up distributing the distances equally in all the buckets while our methods accurately compute the proportions of distances that should go into each bucket. For both datasets, A2 has the same level of errors with those of A1, although the error fluctuates in the spectrum of different bucket width and tends to be larger under smaller bucket width. The reason for this, again, is because A2 works for lower levels of the tree and the number of particles is small.



Fig. 16 Time–Error Product (TEP) of different SDH computation algorithms

Deeper insights on the performance/error tradeoff of different algorithms can help users make justifiable choices. One way to quantify the performance/error tradeoff is the *product of time and error* - an algorithm

with lower *time-error product* (TEP) is obviously preferred. We calculated the TEPs of all tested algorithms and found that, among all settings and algorithms, A4 stands the winner by producing the smallest TEPs under all bucket widths (Fig. 16), although its advantage over A3 is very small in the 8-million atom dataset. Algorithm A3 is only second to A4 with slightly higher TEPs, beating A1 and A2. This clearly shows that A4, although carries a larger error than A3, can still be a viable choice – its performance gain overshadows the loss of accuracy as compared to A3. The gain or loss in time and error may compensate each other in some cases, producing similar TEPs. It is user's choice to pick either A3 or A4. Again, A2 only shows its advantage over A1 under larger w values, indicating that using temporal locality alone is not a viable choice.

Number of simulations: Much time in computation of the first few frames is spent in performing the simulations to update the hash table entries. In our experiments on the dataset of 890K atoms, the number of simulations performed for each frame dropped quickly. In total, 100 frames were processed to compute SDH using algorithm A3. Fig. 17 shows the distribution of simulations performed over 100 frames. We can see that the first frame peaks at 120 simulations. In most of the other frames, no simulations are performed except for few frames for which less than 25 simulations are performed. This clearly states that the hash table utilized in A3 saves running time by reusing the simulations performed in previous frames.



Fig. 17 Number of simulations performed per frame to process 100 frames together under bucket width of 1450

The resolved pairs of cells eliminate direct computation of large number of distances, saving simulation time as well. Fig. 18 shows the number of such distance computations eliminated in SDH processing with different bucket sizes.

Simulation size: The number of points used in every Monte Carlo simulation does not affect the SDH results, as long as sufficient number of points are generated.



Fig. 18 Distances resolved in SDH computation. Datasets: D1 - 890K atoms; D2 - 8 million atoms (y-axis has log scale)

The error shown in Fig. 19 does not change when the number of points in the Monte Carlo simulations goes beyond 50. Thus, our analysis of the Type I error in Section 7.1.2 only gives a loose error bound whereas the actual errors are much lower.



Fig. 19 Effects of simulation size on SDH error

Using noncentral χ^2 distribution: The noncentral χ^2 distribution approximation of the distances between two cells is applied to compare with the Monte-Carlo simulations. Specifically, for each pair of cells, we distribute the distance counts into the relevant buckets based on the values obtained from the Cumulative Distribution Function (CDF) of the noncentral χ^2 distribution. Such values are computed by calling a MATLAB library [33] and cached into a hash table to avoid repeated computations (exactly the same as what we did for the Monte Carlo simulation results). Fig. 20 shows the comparison of errors in the SDH obtained and the running time. The errors generated by using the CDF of noncentral χ^2 are slightly higher than those by the Monte Carlo simulation. This is expected as we know there is a systematic error in using the CDF (Lemma 1) while the Monte Carlo simulations are shown to be very accurate (Section 7.1.2). The simulation-based method also beats the CDF-based method in efficiency. This is because the CDF of noncentral χ^2 distribution has a very complex form [25] therefore the time used for numerical computations in Matlab is non-trivial.



Fig. 20 Histogram errors and computation time using noncentral χ^2 distribution function (CDF) and Monte Carlo simulation (Sim)

Summary: Computation of SDH based on spatial uniformity delivers the significant performance boost over existing algorithm while generating more accurate results. The idea of utilizing the temporal locality can work on top of the spatial uniformity idea to achieve higher performance and also better performance/accuracy tradeoffs. This idea by itself did not show clear advantage, as demonstrated by the bad performance of A2 under small bucket width. Monte Carlo simulation should be the choice in making distance distribution decisions, although the approach based on the CDF of noncentral χ^2 is only marginally worse. The simulationbased approach generates very little error even when the simulation size is small, making it a winner over the CDF-based approach. The advantages of the new algorithm over ADM-SDH become small under large bucket width, but this does not generate a concern since the target of the new algorithm is the smaller bucket width, which is preferred in scientific data analysis.

9.3 Results of GPU Experiments

The GPU versions of the proposed algorithm and ADM-SDH were implemented under CUDA, version 4.0 [34]. The performance of the algorithms was evaluated on an NVIDIA GeForce GTX 570 GPU. We report results for processing the 8-million-atom dataset.

Main results: A comparison of results of different implementations of the proposed algorithm are shown in Fig. 21, in which we show the performance of processing the first frame only using A3. The running time on GPU shows the trend similar to that on CPU, but much faster. The speedup varies with the use of different types of memory. When only the global memory is used, the speedup achieved ranges from 7X to 10.4X.



Fig. 21 Comparing running time and speedup on GPU using different memories. MM: host main memory; GM: GPU global memory; SM: GPU shared memory

The use of shared memory pushes the speedup further by a factor of 2 (i.e., actual speedup ranges from 14.9Xto 25.8X). The speedup is limited by the random memory access patterns emerging due to divergence in the thread computation. The thread divergence also serializes the execution of some of the threads. The size of the tree nodes that are stored in the memory also affect the access patterns due to bank conflicts in shared memory.

The huge speedup under small w values is due to two factors: (1) Parallel processing of the cells in only one level of the density map tree and (2) Reduced divergence in the threads of each GPU block. Even though the computations diverge in processing some pairs of cells, the speedup is achieved by processing on different multiprocessors. Each multiprocessor has its own dedicated shared memory and does not interfere with other multiprocessors' execution.

The separation of simulation and other computations made the algorithm running time almost constant for consecutive frame, for fixed bucket width. Fig. 22 shows the processing time of all 10 frames using the A3 algorithm implemented in both CPU and GPU. Employing the GPUs reduces the computation time of first frame significantly. Also, all the simulations can be done within 100ms, significantly reducing their contribution to the algorithm's running time. Hence, the SDH can be computed efficiently in real time.

Energy efficiency: As a side note, the energy efficiency of the GPU implementations is worth some discussions. Energy consumption has become a major concern in database system design [1]. The product of computation time and active power⁸ consumed for SDH pro-

 $^{^{8}}$ Active power is the power measured for the entire database server less the system idle power. It can be viewed



Fig. 22 Processing time of consecutive frames on GPU. A bucket width of 50 is set for this experiment



Fig. 23 Active energy consumption of CPU and GPU implementations of A3 algorithm under different bucket width

cessing define the energy efficiency of the algorithms. Fig. 23 plots the energy consumed by both CPU and GPU versions of the A3 algorithm. Although the active power consumption of a GPU is a couple of times higher than that of the CPU (46 watts vs. 17 watts as we recorded), the efficiency of the GPU algorithms makes it an energy efficient device for SDH computation - active energy consumption is 5.39 to 9.13 times lower for the GPU code using shared memory. Even for the one that uses only global memory, energy efficiency is 2.51 to 3.81 times higher. To calculate the total energy consumption for the whole machine, we have to add an idle power of 114.5 watts to the active power readings and that will translate into even larger energy savings for the GPU implementations.

GPU implementation of ADM-SDH: We also implemented the ADM-SDH algorithm (A1) on the GPU, and the running time and speedup are shown in Fig. 24. We can see speedup up to 100 times in cases of small w

values, and such performance gain diminishes when wincreases. Recall that, in A1, a density map is chosen for processing such that the cell diagonal length is no greater than w. As w increases, the density map closer to root of the tree is selected. This reduces the computation time in CPU, and in GPU the number of cell pairs processed in parallel decreases. Hence, we see a trend of decreasing speedup achieved by the GPU version. We see a small speedup drop in the GPU algorithm for very small bucket width. This is mainly because the number of cell pairs to be processed increases significantly at the lower level of the tree, increasing the running time. Note that we only show the results of a GPU implementation on global memory, since the utilization of shared memory for ADM-SDH does not further improve the performance. The reason for that is: ADM-SDH processes cells on several levels of the tree (starting from DM_k) therefore we need to pack subtrees instead of single cells into shared memory. However, only a part of each subtree will be used for computation in a random manner thus it is not efficient.



Fig. 24 Comparing running time and speedup of ADM-SDH (A1) on GPU. MM: host main memory (CPU) implementation; GM: GPU global memory (GPU) implementation

Summary: The GPU versions of our algorithm demonstrate the great potential of GPUs in large-scale data analytics. For the SDH problem we tested, speedup over the single-CPU implementation reaches 25X - that is a significant improvement of performance. The speedup decreases under larger bucket width, but it is always the cases of smaller bucket width that make the SDH problem difficult. Such diminish of speedup, as well as the different optimization strategies, however, indicate that GPU programming is a non-trivial task. Finally, the combination of GPU's computing power and efficient algorithm to utilize the spatio-temporal uniformity, delivers very high performance. As a result, we

as the power used for processing the workload. In our experiments, we use a WattsUp power meter to measure total server power consumption.

are able to analyze scientific simulation data in a real time manner.

10 Conclusions and Future Work

An efficient approximate solution to the spatial distance histogram query is provided in this paper. SDH is one of the very important molecular simulation data analysis queries that is frequently applied to a collection of data frames. We use the point region Quad-tree to organize simulation data as we did in our previous work. However, the algorithm presented in this work provides much higher efficiency and accuracy by taking advantage of the data locality and statistical data distribution properties. The algorithm presented makes it practically feasible to analyze data of large number of frames continuously. Its efficiency and accuracy are supported by mathematical analysis and extensive experimental results. We have also shown that, through experiments, utilizing power of modern GPUs gives very significant improvement in the performance. The scientific data analysis can be performed in real time by using such modern hardware systems.

An immediate work of interest is to extend our algorithm to spatial particle distributions other than the uniform pattern. Another important direction of research would be to study the feasibility of computing general *m*-body correlation functions in scientific databases. Such functions, despite the high scientific value they carry, have not been thoroughly studied due to the lack of efficient computing algorithms. We strongly believe the idea based on spatial uniformity as well as GPU programming can be extended to *m*-body correlation function computation. We are in the process of developing solutions for such problems and hope, with the success of such development, to enter an exciting era of computational science endeavours.

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Appendix

A Introduction to χ^2 test

The chi-square test checks if a population is uniformly distributed over different categories. It essentially tests how close the observed values are to the expected values of the model tested. The observed values are the frequencies in categories/classes of the test sample and the expected values are the frequencies in categories/classes under uniform distribution assumption. The smaller the differences between the observed and expected values, the bigger the chances that the sample of data in question follows the claimed distribution. One constrain for the χ^2 test is that it performs poorly when the expected values of each category is less then 5 [20]. The χ^2 goodness-of-fit test is defined as follows:

Definition 2 The χ^2 test is defined for the following hypothesis:

 H_0 : The data are uniformly distributed over categories.

 ${\cal H}_a :$ The data are not uniformly distributed over categories.

The test statistic (χ^2) is defined through the following equation:

$$\chi^{2} = \sum_{i=1}^{k} \frac{(O_{i} - E_{i})^{2}}{E_{i}}$$

where k is the number of categories, O_i is the observed frequency and E_i is the expected frequency of data in each category.

Another value that the χ^2 test needs is the degrees of freedom value, df. The degrees of freedom value is computed as follows: df = cat - 1, where cat is the number of different categories of data we are considering in the test. Once the chi-square value is computed, one can either look up the chi-square table or use software to compute *p*-value. Lastly, the *p*-value is compared to α , the significance value. If $p > \alpha$, the null hypothesis is not rejected, otherwise, the null hypothesis is rejected. The smaller the significance value α , the stronger the test is, meaning if $p > \alpha$, the probability that the observed distribution is equivalent to the expected one is higher. Most accepted α values in the scientific world are 0.001, 0.01 and 0.05, signifying that the probability of making mistake in accepting the null hypothesis is 0.1%, 1% and 5% respectively.

The validity of χ^2 test of the spatial uniformity is supported by the mutual independence/weak dependence assumption of the distribution of the particles over the localized regions. This may result from the fact that the inter-particle forces take effect only within a critical distance, negligibly short compared with the diameter of the localized regions. Therefore, each particle may be treated as the center of a ball with the radius of the critical distance. The distribution of the balls, hence the particles, are mutually independent over the localized regions.

B Distance Distribution Within and Between Two Unit Squares

If two points are randomly and uniformly taken from the same unit square (i.e., one with lateral length 1), the distribution of the distance between such two points has the following probability density function:

$$f(x) = \begin{cases} 2x(x^2 - 4x + \pi) & 0 \le x \le 1\\ 2x[4\sqrt{x^2 - 1} - (x^2 + 2 - \pi) & \\ -4\tan^{-1}\sqrt{x^2 - 1}] & 1 \le x \le \sqrt{2} \end{cases}$$

For two points uniformly sampled from two adjacent unit squares, the distance has the following distribution function:

$$F(x) = \begin{cases} \frac{2x^3}{3} - \frac{x^4}{4} & 0 \le x \le 1\\ \frac{3x^2}{2} - \frac{4x^3}{3} + \frac{x^4}{2} + 2x^2 \arccos(1/x) \\ -\frac{1}{4} - \frac{2(1+2x^2)(x^2-1)^{\frac{1}{2}}}{3} & 1 \le x \le \sqrt{2} \end{cases}$$

$$F(x) = \begin{cases} \frac{2x^2 \arcsin(1/x) - \frac{11}{12} - \frac{x^2}{2} - \frac{4x^3}{3} \\ + \frac{2(1+2x^2)(x^2-1)^{\frac{1}{2}}}{3} & \sqrt{2} \le x \le 2 \end{cases}$$

$$\frac{2(1+2x^2)(x^2-1)^{\frac{1}{2}}}{3} - \frac{75}{12} - \frac{9x^2}{2} \\ + \frac{2(2+x^2)(x^2-4)^{\frac{1}{2}}}{3} + \frac{5x^4}{12} \\ + 2x^2 \arcsin(1/x) \\ + 2x^2 \arccos(2/x) \left[-1 \\ + \frac{5-x^2}{(x^2-4)^{\frac{1}{2}}} \right] & 2 \le x \le \sqrt{5} \end{cases}$$

C Total Number of Simulations Performed

The density map is organized as a grid of $M = n \times n$ cells. We represent the position of each cell by an ordered pair (x, y), where x and y are the horizontal and vertical displacements respectively, of the cell from the top-left corner of the density map. A cell C with displacements i, j is represented by C(i, j). The width or side of each cell is denoted by t (see Fig. 25). We discuss the number of Monte Carlo simulations performed in a density map through a special feature called L-shape (Definition. 3). The number of simulations performed is directly related to the number of distinct L-shapes found in the density map.

Definition 3 *L*-shape of two cells *A* and *B*, L(A, B), is a subset of the density map that includes the two end cells $A(x_A, y_A)$ and $B(x_B, y_B)$ and all the cells with positions

$$(x_A + 1, y_A), (x_A + 2, y_A), \dots, (x_B, y_A)$$
 and
 $(x_B, y_A + 1), (x_B, y_A + 2), \dots, (x_B, y_B - 1)$

or the positions

$$(x_A, y_A + 1), (x_A, y_A + 2), \dots, (x_A, y_B)$$
 and
 $(x_A + 1, y_B), (x_A + 2, y_B), \dots, (x_B - 1, y_B)$



Fig. 25 Illustration of *L*-shape L(A, B) of size d(L(A, B)) = (a, b) in a density map

Without loss of generality we assume $x_A < x_B$ and $y_A < y_B$ in rest of the discussion. It is to be noted that both cells, Aand B, have only one neighbor cell in the L(A, B)-shape.

Definition 4 The size of an L(A, B) shape, which is denoted as d(L(A, B)), is the ordered pair (a, b) where a is the horizontal distance (counted in number of cells) and b is the vertical distance between the cells A and B.

Definition 5 Equivalent *L*-shapes: Let L(A, B) and L(C, D) be two *L*-shapes with sizes d(L(A, B)) = (a, b) and d(L(C, D)) = (c, d). Then L(A, B) is equivalent to L(C, D) (i.e., $L(A, B) \equiv L(C, D)$) iff (a = c and b = d) or (a = d and b = c).

Lemma 2 $L(A, B) \equiv L(C, D)$ iff the minimum and maximum distances between A, B and between C, D are equal. In other words, $L(A, B) \equiv L(C, D)$ iff $dist_{min,max}(A, B) = dist_{min,max}(C, D)$.

Proof Consider two *L*-shapes, L(A, B) and L(C, D) with sizes d(L(A, B)) = (a, b) and d(L(C, D)) = (c, d).

If $L(A, B) \equiv L(C, D)$ then, by the definition 5, d(L(A, B)) = d(L(C, D)). Thus, a = c and b = d or a = d and b = c.

Fig. 25 shows maximum distance between cells A and B.

$$\begin{aligned} dist_{max}(A,B) &= \sqrt{((a+1)*t)^2 + ((b+1)*t)^2} \\ &= \sqrt{((c+1)*t)^2 + ((d+1)*t)^2} \\ &= dist_{max}(C,D) \end{aligned}$$

Similarly for the minimum distance between cells A and B,

$$\begin{aligned} dist_{min}(A,B) &= \sqrt{((a-1)*t)^2 + ((b-1)*t)^2} \\ &= \sqrt{((c-1)*t)^2 + ((d-1)*t)^2} \\ &= dist_{min}(C,D). \end{aligned}$$

Let two pairs of cell (A, B) and (C, D) have same minimum and maximum distance between them i.e.,

$$dist_{min,max}(A,B) = dist_{min,max}(C,D)$$

or in an equivalent form:

$$\sqrt{((a-1)*t)^2 + ((b-1)*t)^2} = \sqrt{((c-1)*t)^2 + ((d-1)*t)^2}$$

Table 2 Number of L-shapes for each value of a

a	0	1	2	3	 n-2	n-1
# L-shapes	n-1	n-1	n-2	n-3	 2	1

The equation holds only if (a = c and b = d) or (a = d and b = c). Thus, $d(L(A, B)) \equiv d(L(C, D))$. By definition, if two *L*-shapes have same size, they are equivalent. \Box

Theorem 2 The number of distinct L-shapes (regardless of position) in a density map with $M = n^2$ cells is $\frac{n(n+1)}{2} - 1$.

Proof The form of each L-shape L(A, B) is defined by its size d(L(A, B)) = (a, b), where $0 \le a \le n - 1$ and $0 \le b \le n - 1$. But, since the L shapes with size (a, b) are equivalent to the L-shapes with size (b, a) we need only to count the L-shapes with size (a, b) where $b \ge a$ and $b \ne 0$. The number of such L-shapes for given values of $a = 1, 2, \ldots n - 1$ are $n - 1, n - 2 \ldots, 1$ respectively. For a = 0 there are n - 1 L-shapes. Obviously, the total number of all distinct L-shapes of size (a, b) is $\frac{n*(n+1)}{2} - 1$.

As the number of distinct Monte Carlo simulations performed in an RDM is equal to the number of distinct Lshapes, the total number of simulation performed to compute SDH is bound by O(M).